approval from OMB to conduct customer surveys over the next three

As part of a comprehensive program, the National Center for Health Statistics (NCHS) plans to continue to assess its customers' satisfaction with the content, quality and relevance of the information it produces. NCHS will conduct voluntary customer surveys to assess strengths in agency products and services and to evaluate how well it addresses the emerging needs of its data users. Results of these surveys will be used in future planning initiatives.

The data will be collected using a

The data will be collected using a combination of methodologies appropriate to each survey. These may

include: Evaluation forms, mail surveys, focus groups, automated and electronic technology (e.g., e-mail, Web-based surveys), and telephone surveys. Systematic surveys of several groups will be folded into the program. Among these are Federal customers and policy makers, state and local officials who rely on NCHS data, the broader educational, research, and public health community, and other data users. Respondents may include data users who register for and/or attend NCHS sponsored conferences; persons who access the NCHS Web site and the detailed data available through it; consultants; and others. Respondent data items may include (in broad

categories) information regarding respondent's gender, age, occupation, affiliation, location, etc., to be used to characterize responses only. Other questions will attempt to obtain information that will characterize the respondents' familiarity with and use of NCHS data, their assessment of data content and usefulness, general satisfaction with available services and products, and suggestions for improvement of surveys, services and products.

The resulting information will be for NCHS internal use. There is no cost to respondents other than their time to participate in the survey.

### ESTIMATED ANNUALIZED BURDEN TABLE

Type of survey	Respondents	Number of respondents	Number of responses/ respondent	Average bur- den/response (in hours)	Total burden hours
Questionnaire for conference registrants/attendees.	Public/private researchers, Consultants, and others.	3,000	1	10/60	500
Focus groups	Public/private researchers, Consultants, and others.	240	1	1	240
Web-based	Public/private researchers, Consultants, and others.	3,600	1	10/60	600
Other customer surveys	Public/private researchers, Consultants, and others.	1,200	1	15/60	300
Total		8,040			1,640

Dated: October 6, 2010.

#### Carol E. Walker.

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

[FR Doc. 2010–25694 Filed 10–12–10; 8:45 am]

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# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

[60Day-11-0776]

### Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, call 404–639–5960 or send comments to Carol E. Walker, CDC Acting Reports Clearance Officer, 1600

Clifton Road, MS–D74, Atlanta, GA 30333 or send an e-mail to *omb@cdc.gov*.

Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Written comments should be received within 60 days of this notice.

## **Proposed Project**

Economic Analysis of the National Breast and Cervical Cancer Early Detection Program—Revision—Division of Cancer Prevention and Control, National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC). Background and Brief Description

CDC administers the National Breast and Cervical Cancer Early Detection Program (NBCCEDP), the largest organized cancer screening program in the United States. The NBCCEDP provides critical breast and cervical cancer screening services to uninsured and underserved low-income women in all 50 States, the District of Columbia. five U.S. territories, and 12 American Indian/Alaska Native organizations. The program provides breast and cervical cancer screening for eligible women who participate in the program as well as diagnostic procedures for women who have abnormal findings. During the past decade, the NBCCEDP has provided over 9.2 million breast and cervical cancer screening and diagnostic exams to over 3.7 million low-income women. Those who are diagnosed with cancer through the program are eligible for Medicaid coverage through the Breast and Cervical Cancer Prevention and Treatment Act passed by Congress in 2000.

In 2008, CDC received OMB approval to collect one year of activity-based economic cost data from NBCCEDP grantees. In 2009, CDC received OMB approval to collect two additional cycles of cost data for fiscal year 2009 (FY09)

and fiscal year 2010 (FY10) (OMB No. 0920–0776, exp. 03/31/2011). Respondents are the 68 programs participating in the NBCCEDP. Information is collected through a webbased Cost Assessment Tool (CAT) and includes: Staff and consultant salaries, screening costs, contracts and material costs, provider payments, in-kind contributions, administrative costs, allocation of funds and staff time devoted to specific program activities.

CDC requests OMB approval for a sixmonth extension of the current approval period in order to complete the third year of data collection. Based on our experience with previous data collection cycles, 20 grantees (30% of the total 68 grantees) will not be able to meet the current data collection deadline of 3/31/2011. These programs will complete their fiscal year (FY) closeout process in April or May 2011.

As a result, these programs will not be prepared to submit data to CDC until their FY is complete and records have been reconciled. The requested sixmonth extension period will provide the time they need to complete their closeout process and conduct data quality checks before submitting information to CDC. The requested sixmonth extension will improve the quality and completeness of information used for planned data analysis, and ensure CDC's authority to receive late submissions.

The activity-based cost data will be used to evaluate grantees to ensure the most appropriate use of limited program resources in delivering program services such as screening, diagnostic services, case management and outreach. The detailed cost data will allow CDC to determine the costs of various program components, identify factors that impact

average cost, perform cost-effectiveness analysis and budget impact analysis of the program, and allocate program resources more effectively and efficiently. The collection of economic cost information complements the measures of NBCCEDP effectiveness collected as Minimum Data Elements (0920–0571, exp. 11/30/2012).

In this Revision request, there are no proposed changes to the data collection instrument, data collection methodology, or the estimated burden per response. The only changes are a decrease in the estimated number of respondents (the number of late responders) and a six-month extension of the data collection period. All information is collected electronically. There are no costs to respondents other than their time.

### ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Number of respondents	Number of responses per respondent	Average burden (in hrs)	Total burden (in hrs)
NBCCEDP grantee	20	1	22	440

Dated: October 6, 2010. Carol E. Walker,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## Health Resources and Services Administration

## Agency Information Collection Activities: Submission for OMB Review; Comment Request

Periodically, the Health Resources and Services Administration (HRSA) publishes abstracts of information collection requests under review by the Office of Management and Budget (OMB), in compliance with the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35). To request a copy of the clearance requests submitted to OMB for review, e-mail paperwork@hrsa.gov or call the HRSA Reports Clearance Office on (301) 443—1129.

The following request has been submitted to the Office of Management and Budget for review under the Paperwork Reduction Act of 1995: Proposed Project: Healthcare Integrity and Protection Data Bank for Final Adverse Information on Health Care Providers, Suppliers and Practitioners (45 CFR 61)—OMB No. 0915–0239— Revision

This is a request for revision and extension of OMB approval of the information collections contained in regulations found in 45 CFR Part 61 governing the Healthcare Integrity and Protection Data Bank (HIPDB) and the forms to be used in reporting information to and requesting information from the HIPDB cleared under OMB No. 0915-0239. An additional form entitled, "Instructions for Registering as an NPDB-HIPDB Self-Querier," has been included to meet identity proofing and e-authentication requirements stipulated in the *E*– Authentication Guidance for Federal Agencies (OMB M-04-04) and National Institutes of Standards and Technology's (NIST) Draft Special Publication 800–63–1, *Electronic* Authentication Guidelines. The burden estimate for self-queriers has been adjusted from the original OMB approval to reflect this new registration process. The HIPDB is authorized by section 1128E of the Social Security Act (hereinafter referred to as section 1128E), as added by section 221(a) of the Health Insurance Portability and

Accountability Act of 1996. Section 1128E directs the Secretary of Health and Human Services (the Secretary) to establish a national health care fraud and abuse data collection program for the reporting and disclosing of certain final adverse actions (excluding settlements in which no findings of liability have been made) taken against health care providers, suppliers, or practitioners. It also directs the Secretary to maintain a database of final adverse actions taken against health care providers, suppliers, or practitioners. The regulations implementing section 1128E governing the operation of the HIPDB are codified at 45 CFR Part 61. The HIPDB became operational November 22, 1999.

Approval is requested to continue the following reporting data collection and disclosure requirements and the ensuing HIPDB forms along with the instructions. The recordkeeping, reporting, and disclosure requirements are specified in the regulations to implement the HIPDB. Numbers in the table may not add up exactly due to rounding. *Please note* the burden for Administrative Forms has been accounted for in the NPDB OMB clearance renewal submission.

The annual estimate of burden is as follows: